Orotidine-5'-phosphate decarboxylase gene, gene construct comprising this gene and its use

5 The invention relates to an orotidine-5'-phosphate decarboxylase gene having the sequence SEQ ID No. 1 or its homologs, to a gene construct comprising this gene or its homologs, and to its use. The invention additionally relates to vectors or organisms comprising an orotidine-5'-phosphate decarboxylase gene having 10 the sequence SEQ ID No. 1 or its homologs.

The invention further relates to a process for producing uracil-auxotrophic microorganisms and to a process for inserting DNA into uracil-auxotrophic microorganisms.

Vitamin B2, also called riboflavin, is essential for humans and animals. Vitamin B2 deficiency is associated with inflammations of the mucosa of the mouth and throat, pruritus and inflammations in skin folds and similar cutaneous lesions, conjunctival 20 inflammations, reduced visual acuity and clouding of the cornea. In babies and children, cessation of growth and weight loss may occur. Vitamin B2 therefore has economic importance in particular as vitamin supplement in cases of vitamin deficiency and as animal feed supplement. It is additionally used as food color, for example in mayonnaise, ice cream, blancmange etc..

Vitamin B2 is prepared either chemically or microbially (see, for example, Kurth et al., 1996, Riboflavin, in: Ullmann's Encyclopedia of industrial chemistry, VCH Weinheim). In the chemical preparation processes, riboflavin is usually obtained as pure final product in multistage processes, it being necessary to employ relatively costly starting materials such as, for example, D-ribose. An alternative to the chemical synthesis of riboflavin is the preparation of this substance by microorganisms. The starting materials used in this case are renewable raw materials such as sugars or vegetable oils. The preparation of riboflavin by fermentation of fungi such as Eremothecium ashbyii or Ashbya gossypii is known (The Merck Index, Windholz et al., eds. Merck & Co., page 1183, 1983), but yeasts such as, for example, Candida, Pichia and Saccharomyces or bacteria such as, for example, 40 Bacillus, clostridia or corynebacteria have also been described as riboflavin producers.

DE 44 20 785 describes six riboflavin biosynthesis genes from Ashbya gossypii, and microorganisms which have been transformed 45 with these genes, and the use of such microorganisms for riboflavin synthesis.

Cowerze . czewo

To date, genes have been inserted into fungal riboflavin producers such as Ashbya gossypii via the markers leu2 (leucine auxotrophy), thr4 (threonine auxotrophy) or kan (kanamycin resistance) (WO 92/00379). A further marker described in yeasts is met15 (methionine auxotrophy, Cost et al., Yeast, Vol. 12, 1996: 939 - 941). The disadvantage of this marker is that either the transformation efficiency is very low and/or antibiotics must be continuously added for the selection. However, in each case, counterselection for loss of the marker with retention of the inserted genes in microorganisms is impossible or possible only with very great effort, so that it is usually no longer possible to insert further genes with these markers into the microorganisms. It is therefore desirable to have a selection marker which displays high transformation efficiency, is easily

15 selectable and makes counterselection possible.

Genet., Vol. 27, 1995: 536 - 540).

The orotidine-5'-phosphate decarboxylase gene (= URA3 gene) from Saccharomyces cerevisiae is one of the classical markers having the required properties and usable for transforming genes into 20 microorganisms such as yeasts and fungi. The isolation of species-specific URA3 genes and the isolation of the corresponding gene from fungi (= pyrG) and the sequences thereof from Pichia stipitis, Candida boidinii, Kluyveromyces marxianus, Yamadazyma ohmeri, Candida maltosa, Aspergillus niger, Aspergillus oryzae, Aspergillus nidulans, Mucor circinelloides, Phycomyces blakesleeanus, Penicillium chrysogenum, and Aspergillus awamori have been described in a number of studies (Appl. Environ. Microbiol., Vol. 60, No. 12, 1994: 4245 - 4254, Nucl. Acids Res., Vol. 18, No. 23, 1990: 7183, J. Ferment. Bioeng., Vol. 73, No 4, 1992: 255 - 260, Yeast, Vol. 9, 1993: 30 677 - 681, Yeast, Vol. 10, 1994: 1601 - 1612, Curr. Genet., Vol. 23, 1993: 205 - 210, Nucl. Acids Res., Vol.16, No. 5, 1988: 2339, Curr. Genet., Vol. 16, 1989: 159 - 163, Gene, Vol. 61, 1987: 385 - 399, Gene, Vol. 116, 1992: 59 - 67, Mol. Gen. Genet., Vol. 224, 1990: 269 - 278, Nucl. Acids Res., Vol. 16, No. 16, 1988: 35 8177, Nucl. Acids Res., Vol. 18, No. 23, 1990: 7183 and Curr.

Studies by Rose et al. (Gene, Vol. 29, 1984: 113 - 124) have shown that the URA3 gene from Saccharomyces cerevisiae is in fact 40 capable of complementation of a corresponding mutation (pyrF gene = URA3) in prokaryotes such as Escherichia coli, and can be useful as selection marker.

However, genetic studies on riboflavin synthesis by Ashbya
45 gossypii (vitamin B2 synthesis) have shown that the URA3 gene
from Saccharomyces derevisiae or the pyrF gene from Escherichia
coli are [sic] not capable of complementation of

uracil-auxotrophic Ashbya gossypii mutants, and therefore these genes cannot be used for cloning genes into Ashbya gossypii.

Attempts have therefore been made, because that [sic] gene from 5 Ashbya gossypii corresponding to the URA3 gene or pyrF gene is unknown, to clone it. Attempts at cloning the Ashbya gene by the methods described in the literature via, for example, hybridization with URA3 gene fragments or via degenerate oligonucleotides based on conserved amino-acid sequences of various orotidine-5'-phosphate decarboxylases and screening a cDNA library using these oligonucleotides and the PCR technique were unsuccessful (Bergkamp et al. Yeast, Vol. 9, 1993: 677 - 681, Piredda et al., Yeast, Vol. 10, 1994: 1601 - 1612, Benito et al., Gene, Vol. 116, 1992: 59 - 67 and Diaz-Minguez et al., Mol. 15 Gen. Genet., Vol. 224, 1990: 269 - 278).

It is an object of the present invention therefore to provide an easily selectable marker which can be transformed with high yield and is easily counterselectable and which makes it possible to insert genes into microorganisms.

We have found that this object is achieved by the novel orotidine-5' phosphate decarboxylase [lacuna] having the sequence SEQ ID NO: 1 or its homologs which have at least 80% homology with the sequence SEQ ID NO: 1.

Homologs of the novel orotidine-5 -phosphate decarboxylase gene having the sequence SEQ ID NO: 1 mean, for example, allelic variants which have at least 80% homology at the derived amino-acid level, preferably at least 90% homology, very particularly preferably at least 95% homology. The amino-acid sequence derived from SEQ ID NO: 1 is to be seen in SEQ ID NO: 1. Allelic variants comprise, in particular, functional variants which are obtainable by deletion, insertion or substitution of nucleotides from the sequence depicted in SEQ ID NO: 1, the intention being, however, that the enzymatic activity of the

intention being, however, that the enzymatic activity of the derived synthesized proteins advantageously be retained for the insertion of one or more genes. However, if the intention is to produce mutants in the orotidine-5'-phosphate decarboxylase gene with the aid of SEQ ID NO: 1 and its homologs in the novel

process for producing uracil-auxotrophic microorganisms, non-functional genes will be used, that is to say genes which lead to enzymatically inactive proteins. In this case, it is advantageous to use sequences which display homologies with SEQ ID NO: 1 or its homologs advantageously at the 3' and 5' ends.

acelore of the seco

20

104. DIL

Homologs of SEQ ID NO: 1 additionally mean, for example, fungal or plant homologs, truncated sequences, single-stranded DNA or RNA of the coding and noncoding DNA sequence. Homologs of SEQ ID NO: 1 have at the DNA level a homology of at least 60%, preferably of at least 70%, particularly preferably of at least 80%, very particularly preferably of at least 90%, over the complete DNA region indicated in SEQ ID NO: 1.

Homologs of SEQ ID NO: 1 also mean derivatives such as, for

10 example, promoter variants. The promoters upstream of the
indicated nucleotide sequences may be modified by one or more
nucleotide exchanges, by insertion(s) and/or deletion(s) without,
however, the functionality or activity of the promoters being
impaired. It is additionally possible for the promoters to have
their activity increased by modifying their sequence, or to be
completely replaced by more active promoters even from
heterologous organisms.

Derivatives also mean variants whose nucleotide sequence in the 20 region from -1 to -200 in front of the start codon have [sic] been modified so as to alter, preferably increase, gene expression and/or protein expression.

It is possible and preferred for SEQ ID NO: 1 or its homologs to 25 be isolated from microorganisms of the family Metschnikowiaceae, particularly preferably from microorganisms of the genera Eremothecium, Ashbya or Nematospora, very particularly preferably from microorganisms of the genus and species Eremothecium ashbyii or Ashbya gossypii.

 $>_{ exttt{The novel gene construct means the URA3 gene sequences [sic] SEQ}$ ID No. 1 and its homologs which have been functionally linked to one or more regulatory signals, advantageously to increase gene expression. Examples of these regulatory sequences are sequences 35 to which inducers or repressors bind and thus regulate the expression of the nucleic acid. In addition to these novel regulatory sequences the natural regulation of these sequences in front of the actual structural genes can still be present and, where appropriate, have been genetically modified so that the 40 natural regulation has been switched off and the expression of the genes has been increased. The gene construct can, however, also have a simpler structure, that is to say no additional regulatory signals have been inserted in front of the sequence SEQ ID No. 1 or its homologs, and the natural promoter with its 45 regulation has not been deleted. Instead, the natural regulatory sequence has been mutated so that regulation no longer takes

place, and gene expression is enhanced. The gene construct may

construct.

additionally advantageously comprise one or more so-called enhancer sequences functionally linked to the promoter and making increased expression of the nucleic acid sequence possible. It is also possible to insert at the 3' end of the DNA sequences 5 additional advantageous sequences, such as further regulatory elements or terminators. The URA3 genes may be present in one or more copies in the gene construct, and the gene or genes can also be inactivated. It is possible with the aid of this or these inactivated genes to generate uracil-auxotrophic mutants in the 10 novel process. It is advantageous for further genes to be present in the gene construct for insertion of further genes into a microorganism. These genes may be located inside a URA3 gene, in which case there ought advantageously to be an intact copy of the URA3 gene and/or another selectable gene such as leu2, thr4 or 15 kan present in the construct, or they can be located outside the URA3 gene. Even if an intact URA3 gene is present in the construct, further markers such as those mentioned above can, where appropriate, be present for selection in the gene

Advantageous regulatory sequences for the novel process are present, for example, in promoters such as cos, tac, trp, tet, trp-tet, lpp, lac, lpp-lac, lacIq, T7, T5, T3, gal, trc, ara, SP6, λ-P_R or λ-P_L promoter and are advantageously used in Gram-negative bacteria. Further advantageous regulatory sequences are present, for example, in the Gram-positive promoters amy and SPO2, in the yeast or fungal promoters ADC1, MFα, AC, P-60, CYC1, GAPDH, TEF, rp28, ADH or in the plant promoters CaMV/35s, SSU, OCS, lib4, usp, STLS1, B33, nos or in the ubiquitin or phaseolin promoter. Also advantageous in this connection are the promoters of pyruvate decarboxylase and of methanol oxidase from, for example, Hansenula. It is also possible to use artificial promoters for the regulation.

It is possible in principle to use all natural promoters with their regulatory sequences like those mentioned above for the novel process. It is also possible and advantageous in addition to use synthetic promoters.

The gene construct may, as described above, also comprise further genes which are to be inserted into the microorganisms. These genes can be inserted inside or outside the marker genes such as ura3, leu2, thr4 or kan. It is possible in principle for all types of genes to be inserted into microorganisms with the aid of the novel URA3 gene having the sequence SEQ ID NO: 1 or its homologs. It is possible and advantageous to insert and express in host organisms regulatory genes such as genes for inducers, repressors or enzymes which intervene by their enzymatic activity

in the regulation, or one or more or all genes of a biosynthetic pathway such as the genes of riboflavin biosynthesis such as, for example, the rib genes or genes of biosynthetic pathways which lead to other fine chemicals, secondary metabolites or proteins, such as the genes of biotin, lysine, methionine, vitamin B12 or carotenoid biosynthesis, or genes which lead to flavorings, growth promoters or odoriferous substances, or individual genes for enzymes such as proteases or lipases, via the URA3 sequence. These genes can be heterologous or homologous in origin. The inserted genes may have their own promoter or else be under the control of the promoter of the sequence SEQ ID No. 1 or its homologs.

For expression in the abovementioned host organism, the gene 15 construct is advantageously inserted into a vector such as, for example, a plasmid, a phage or other DNA, which makes optimal expression of the genes in the host possible. Examples of suitable plasmids are, in E. coli, pLG338, pACYC184, pBR322, puc18, puc19, pkc30, pRep4, pHS1, pHS2, pPLc236, pMBL24, pLG200, 20 pUR290, pIN-III¹¹³-B1, λgtl1 or pBdCI, in Streptomyces, pIJ101, pIJ364, pIJ702 or pIJ361, in Bacillus, pUB110, pC194 or pBD214, in Corynebacterium, pSA77 or pAJ667, in fungi, pALS1, pIL2 or pBB116, in yeasts, 2µM, pAG-1, YEp6, YEp13 or pEMBLYe23, or, in plants, pLGV23, pGHlac+, pBIN19, pAK2004 or pDH51. Said plasmids represent a small selection from the possible plasmids. Further plasmids are well known to the skilled worker and can be found, for example, in the book Cloning Vectors (Eds. Pouwels P. H. et al. Elsevier, Amsterdam-New York-Oxford, 1985, ISBN 0 444 904018).

The gene construct advantageously comprises, for expression of the other genes present, additionally 3' and/or 5' terminal regulatory sequences to enhance expression, which are selected for optimal expression depending on the selected host organism and gene or genes.

35

These regulatory sequences are intended to make specific expression of the genes and of the [sic] protein expression possible. This may mean, depending on the host organism, for example that the gene is expressed or overexpressed only after induction, or that it is immediately expressed and/or overexpressed.

The regulatory sequences or factors may moreover preferably have a beneficial effect on expression of the introduced genes, and thus increase it. It is possible in this way for the regulatory elements to be enhanced advantageously at the transcription level by using strong transcription signals such as promoters and/or

enhancers. However, in addition, it is also possible to enhance translation by, for example, improving the stability of the mRNA.

In a further embodiment of the vector, the novel gene construct 5 can also be advantageously introduced in the form of a linear DNA into the microorganisms and be integrated into the genome of the host organism by heterologous or homologous recombination. This linear DNA can consist of a linearized plasmid or only of the gene construct as vector.

10

Host organisms suitable in principle for the novel gene construct are all prokaryotic or eukaryotic organisms. The host organisms advantageously used are microorganisms such as bacteria, fungi, yeasts, animal or plant cells. Fungi or yeasts are preferably used, particularly preferably fungi, very particularly preferably fungi of the family Metschnikowiaceae such as Eremothecium, Ashbya or Nematospora.

The invention additionally relates to a process for producing uracil-auxotrophic microorganisms. To generate uracil-auxotrophic mutants, the orotidine-5'-phosphate decarboxylase gene having SEQ ID NO: 1 or its homologs are modified, for example by mutagenesis, in such a way that the protein encoded by the gene is inactivated. This inactivated gene is subsequently introduced into a microorganism, for example by transformation or electroporation. Finally, homologous recombination in the microorganisms results in auxotrophic mutants which can be screened via their resistance to 5-fluoroorotic acid (see Boeke et al., Mol. Gen. Genet., Vol. 197, 1984: 345 - 346).

201 30

The invention further relates to a process for inserting DNA into organisms, which comprises inserting into an organism, preferably a microorganism, which is deficient in an orotidine-5'-phosphate decarboxylase gene (= URA3 gene) a vector which comprises an intact URA3 gene having the sequence SEQ ID NO: 1 or its homologs, advantageously together with further DNA, preferably with at least one other gene, and cultivating this organism on or in a culture medium which contains no uracil. Only these organisms which have acquired the vector are able to grow in this medium. A linear DNA is preferably used as vector in this process. The microorganisms preferably used in this process are fungi, especially of the family Metschnikowiaceae such as Eremothecium, Ashbya or Nematosprora [sic], particularly preferably microorganisms of the genus Ashbya.

45 It is also possible to use as vector any suitable plasmid (but especially a plasmid which harbors the origin of replication of the 2m plasmid from S. cerevisiae) which undergoes autonomous

replication in the cell, but also, as described above, a linear DNA fragment which is integrated into the genome of the host. This integration can take place by heterologous or homologous recombination. But preferably, as mentioned, by homologous recombination (Steiner et al., Genetics, Vol. 140, 1995: 973 - 987).

The novel URA3 gene having the sequence SEQ ID NO: 1 or its homologs can advantageously be used as selection markers in the novel process. It is possible and preferred to insert genes using this selection marker genes [sic] into Ashbya gossypii.

An additional advantage is that on transformation of Ashbya gossypii it is possible to select with the aid of this gene, 15 without the need to use foreign DNA (i.e. DNA not derived from Ashbya gossypii).

It is possible on transformation of Ashbya gossypii with the gene having SEQ ID NO: 1 or its homologs also to insert any other genes. This makes it possible to construct strains which harbor single genes or a plurality of genes in several copies either on plasmids or in the genome.

It is further possible to construct Ashbya strains in which 25 chromosomal copies of genes have been destroyed by the insertion of the URA3 gene having SEQ ID NO: 1 or its homologs.

A particular advantage of the AgURA3 gene is the possibility of using the marker several times in succession in the same strain.

If identical nucleotide sequences are placed 5' and 3' of the gene in the same orientation (so-called direct repeats), it is possible if required to delete the Aguras marker again by homologous recombination and selection on uracil— and

35 FOA-containing medium, and then in another round insert additional DNA with the aid of this gene. Another advantage is the distinctly greater transformation efficiency by comparison with the markers thr, leu or kan.

In the novel process, the vector comprises as other gene at least 40 one gene of riboflavin synthesis. Genes of riboflavin synthesis mean those genes which are involved in synthesis in the entire metabolism of riboflavin producers such as Ashbya.

Examples:

Example 1:

5 Production of a genomic gene bank from Ashbya gossypii ATCC10895

Genomic DNA from Ashbya gossypii ATCC10895 was prepared by the process described in WO97/03208. The genomic gene bank derived from this DNA was constructed in pRS314 and in YEp351 (Hill et al., Yeast, Vol. 2, 1986: 163 - 167) by the method described in Sambrook, J. et al. (1989) Molecular cloning: A laboratory manual, Cold Spring Harbor Laboratory Press or in [lacuna] F.M. et al. (1994) Current protocols in molecular biology, John Wiley and Sons. As can be inferred from, for example, WO97/03208, other plasmids, such as plasmids of the pRS series (Sikorski and Hieter, Genetics, 1989: 19-27) or cosmids such as, for example, SuperCosl (Stratagene, La Jolla, USA), are also suitable for producing the gene bank.

20 Example 2:

It was initially attempted to clone the gene for the orotidine-5'-phosphate decarboxylase (= OMP-DC) from Ashbya gossypii via functional complementation of a corresponding URA3-auxotrophic mutant of Saccharomyces cerevisiae.

To this end, a gene bank was constructed from genomic Ashbya gossypii DNA in pRS314 (as described in Example 1). This DNA was used to transform the S. cerevisiae strain MW3317-21A (genotype:

30 MAT α, trp1, ade8ΔKpn, ura3-52, hom3-10, met13, met4, ade2, his3-Kpn, see, for example, Kramer et al., Mol. Cell. Biol. 9, 1989: 4432-4440), by the lithium acetate method (see, for example, Kramer et al., Mol. Cell. Biol. 9, 1989: 4432-4440). No clone in which the genomic deletion of the ura3 gene of the S.

35 cerevisiae strain was complemented by a gene fragment from Ashbya was obtained.

The attempt to clone the URA3 gene of Ashbya gossypii via functional complementation in a pyrF mutant of E. coli also 40 failed.

Example 3:

An attempt to clone the OMP-DC gene from Ashbya gossypii by hybridization with a fragment of the corresponding gene from 45 Saccharomyces cerevisiae was also unsuccessful.

For this purpose, the complete URA3 gene from Saccharomyces cerevisiae (gene bank entry yscodcd) was used as probe (length 1.1 kb) in order to screen a genomic cosmid gene bank from Ashbya gossypii (see Example 1). The experiment was carried out as described in Sambrook, J. et al. (1989) Molecular cloning: A laboratory manual, Cold Spring Harbor Laboratory Press or Ausubel, F.M. et al. (1994) Current protocols in molecular biology, John Wiley and Sons, using hybridization temperatures of 52°C to 68°C. It was not possible to identify in the gene bank any

10 clones which provided a positive signal with the URA3 gene from S. cerevisiae as probe.

Example 4:

15 In the next approach, it was attempted to clone the gene for OMP-DC from Ashbya gossypii by amplification of gene fragments using degenerate oligonucleotides and the PCR technique.

For this experiment, the known amino-acid sequences of the 20 various orotidine-5'-phosphate decarboxylases from the following organisms were compared, and regions showing maximum conservation in all the enzymes were selected:

Aspergillus niger (Acc. number: P07817)
Aspergillus nidulans (Acc. number: P10652)
Schizosaccharomyces pombe (Acc. number: P14965)
Penicillium chrysogenum (Acc. number: P09463)
Kluyveromyces lactis (Acc. number: P07922)
Candida albicans (Acc. number: P13649)
Neurospora crassa (Acc. number: P05035)
Ustilago maydis (Acc. number: P15188)
Saccharomyces cerevisiae (Acc. number: P03962)
Drosophila melanogaster (Acc. number: Q01637)
Mouse (Acc. number: P13439)
Human (Acc. number: P11172)

The numbers given in parentheses are derived from the SWISS&PIR-Translated Datenbank Release 103.

40 Degenerate olgonucleotides [sic] were synthesized on the basis of this information.

Degenerate oligonucleotides mean oligonucleotides in which mixtures of nucleotides have been incorporated at several 45 positions during the synthesis.

In this connection, R represents G or A, Y represents C or T, W represents A or T, M represents A or C, K represents G or T, S represents C or G, H represents A, C or T, V represents A, C or G, B represents C, G or T, D represents A, G or T, and N represents G,A,T or C.

The following oligonucleotides were used:

(YEQ ID MO:3)

URA3-A: 5'-XTNGGNCCNTAYATHTGY-3'

URA3-B: 5'-TAYTGYTGNCCNARYTTRTCNCC-3"

URA3-C: 5'-TTYYTNATHTTYGARGAYMGNAARTT-3'

URA3-D: 5'-GCNARNARNARNCCNC-3'

Using these oligonucleotides as primers, PCRs were carried out 15 with genomic DNA from Ashbya gossypii as template.

The following primer combinations were used:

URA3-A + URA3-B; URA3-A + URA3-D; URA3C + URA3-B and URA3-C + 20 URA3-D.

The following hybridization temperatures were used:

52°C, 48°C, 44°C, 40°C and 37°C.

The products resulting from the PCRs were cloned by conventional methods into the vector pGEMT (Promega) and were sequenced. It was not possible to amplify any fragments which showed homology with the known OMP-DC genes mentioned above.

Example 5:

30

35

A cDNA bank was constructed from Ashbya gossypii as described in DE 44 20 785 Al (Example 1).

Example 6:

Analysis of nucleic acid sequences in the gene bank

40 Single clones were selected from E.coli clones which comprised the gene bank from Ashbya gossypii described in Example 5. The cells were cultivated by conventional methods in suitable media (e.g. Luria broth with 100 mg/l ampicillin), and plasmid DNA was isolated from these cells.

Oligonucleotides which hybridize in the vector portion were used as primers for sequencing the cDNA clones. Fragments of the cloned cDNAs were detected in this way. The sequences were analyzed as described in Example 7.

5
Example 7:

A computer-assisted analysis of the nucleotide sequences found was carried out by comparisons of newly identified sequences with 10 existing DNA and protein data banks using the following algorithms, e.g. with BLAST algorithms (Altschul et al. (1990) J. Mol. Biol. 215, 403-410), the Clustal algorithm with the aid of the PAM250 weighting table or the Wilbur-Lipman DNA alignment algorithm (as implemented, for example, in the program package 15 MegAlign 3.06 supplied by DNAstar). It was possible in this way to discover similarities of the newly discovered sequences with previously known sequences, and to describe the function of novel genes or part-sequences of genes.

20 Example 8:

Identification of E. coli clones which harbor the gene for OMP-DC from Ashbya gossypii (AgURA3).

25 After examination of a large number of clones as described in Examples 6 and 7 (> 100 clones), a sequence which showed similarities with known OMP-DC genes was found. This homologous process was then used to screen the genomic Ashbya gene bank (see Example 1) once again, and it was possible to identify clones and cosmids which gave a specific positive signal and harbored a common 1.3 kb XhoI-EcoRI fragment. Sequencing of the clones produced the sequence as described in SEQ ID NO: 1. The sequence shows similarities with known URA3 genes and codes for a protein about 29246 Dalton in size.

Example 9:

Disruption of the chromosomal copy of the AgURA3 gene with antibiotic resistance genes

Disruption of a gene means destruction of the functionality of a genomic copy of the gene either by (a) deleting part of the gene sequence or by (b) of the [sic] interrupting the gene by introducing a piece of foreign DNA into the gene or by (c)

45 replacing part of the gene by foreign DNA. Any foreign DNA can be used, but it is preferably a gene which effects resistance to any

suitable chemical. Any suitable resistance genes can be used to disrupt genes.

To disrupt the Agura3 gene of Ashbya gossypii ATCC10895, the sanamycin resistance gene from Tn903, which [lacuna] under the control of the TEF promoter of Ashbya gossypii (see Yeast 10, pages 1793-1808, 1994 or WO92/00379), was used. The gene is flanked 5' and 3' by several cleavage sites for restriction endonucleases, so that it was possible to construct a cassette 10 which make [sic] possible any desired constructions of gene disruptions using conventional methods of in vitro DNA manipulation.

The internal 370 bp PstI-KpnI fragment of AgURA3 (position 442 - 15 892 in sequence SEQ ID NO: 1) was replaced by a resistance cassette as outline above. The resulting construct was given the name ura3::G418. The resulting plasmid can be replicated after transformation into E.coli. The XhoI-SphI fragment of the construct ura3::G418 (see Figure 1) was purified by agarose gel electrophoresis and subsequent elution of the DNA from the gel (see Proc. Natl. Acad. Sci. USA 76, 615-619, 1979) and employed to transform Ashbya gossypii. Figure 1 shows in depiction A a restriction map of the coding region of the AgURA3 gene and of the 5'- and 3'-untranslated regions (= 5'-UTR and 3'-UTR).

25 Depiction B shows the situation after insertion of the kanamycin resistance cassette described above (= TEF-kanR).

The fragment was transformed into Ashbya gossypii either by protoplast transformation (Gene 109, 99-105, 1991) or else, 30 preferably, by electroporation (BioRad Gene Pulser, conditions: cuvettes with slit widths of 0.4 mm, 1500V, $25\mu F$, 100Ω). The selection of transformed cells took place on G418-containing solid medium (WO 97/03208).

35 Resulting G418-resistant clones were examined by conventional methods of PCR and Southern blot analysis (Sambrook, J. et al. (1989) Molecular cloning: A laboratory manual, Cold Spring Harbor Laboratory Press and in [lacuna] F.M. et al. (1994) Current protocols in molecular biology, John Wiley and Sons) to find 40 whether the genomic copy of the Agura3 gene was in fact destroyed. Clones whose Agura3 gene was destroyed are uracil-auxotrophic and resistant to 1 mg/ml 5'-fluoroorotic acid (FOA).

Example 10:

Disruption of the chromosomal copy of the AgURA3 gene without using antibiotic resistance genes

A particular advantage of the use of URA3 genes is the possibility of selection both for the presence and for the absence of the gene. It is possible to screen with FOA microorganisms which have a functionally inactivated URA3 gene, 10 and by means of selection for uracil prototrophy to select for a functionally active URA3 gene.

To disrupt the genomic copy of the URA3 gene, for the sake of simplicity an internal fragment (= PstI fragment) of the URA3

15 gene was deleted from the coding region of the gene having the sequence SEQ ID NO: 1 (position 442 to 520 in sequence SEQ ID NO: 1). Transformation of Ashbya gossypii with this deleted ura3 fragment was carried out as described in Example 10. In place of deletion of part-regions of the gene, it is also possible in principle to use all other methods for inactivating the gene, such as mutations via insertions, duplications, reversions, replacement of several nucleotides or point mutations. Point mutations are less preferred because reversion thereof is easy.

The transformants were selected through resistance to FOA. In contrast to wild-type clones, clones which harbor a disruption of the Agura3 gene are resistant to 1 mg/ml FOA.

30 Example 11:

Use of the Agura3 gene for inserting further DNA into A. gossypii.

35 The isocitrate lyase gene described in WO 97/03208 was inserted with the aid of the plasmid pAG100, as described in WO 97/03208 (Example 4 and 5), into AgURA3 disruption mutants of A. gossypii (see Example 9 and 10), using as selection marker in A. gossypii the AgURA3 gene in place of the G418 resistance described.

SEQUENCE LISTING

MB)

(1) GENERAL INFORMATION:

- (i) APPLICANT:
 - (A) NAME: BASF Aktiengesellschaft
 - (B) STREET: Carl Bosch Strasse
 - (C) CITY: Ludwigshafen
 - (D) FEDERAL STATE: Rheinland-Pfalz
 - (E) COUNTRY: G∉rmany
 - (F) POSTAL CODE: D-67056
- (ii) TITLE OF APPLICATION: Orotidine-5'-phosphate decarboxylase gene, gene construct comprising this gene and its use
- (iii) NUMBER OF SEQUENCES: 2
 - (iv) COMPUTER-READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.25 (EPO)
- (2) INFORMATION FOR SEQ D NO: 1:
 - (i) SEQUENCE /CHARACTERISTICS:
 - (A) /LENGTH: 1380 base pairs
 - (B) TYPE: nucleic acid
 - (C √ STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA (genomic)
 - (iii) HYPOTHETICAL: NO
 - (iii) [sic] ANTISENSE: NO
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Ashbya gossypii
 - (vii) IMMEDIATE SOURCE:
 - (B) CLONE: ura3
 - (ix) FEATURES:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: 210..1013

							10		1							
	(:	ix)	F	EATU					/							
						NAME										
				(B)]	LOCAT	NOL	: 1/	. 199							
	, .	: \	E.	וויית מ	RES:											
	(.	ix)	F.			NAME	/KEY	: b"	UTR							
				•		LOCA				1380						
				•	•			/								
	(:	xi)	S	EQUE	NCE	DESC	RIPT	TON:	SEQ	ID	NO:	1:				
CTCG.	AGCA	AC I	CATT	GGAA	G CC	CTTC	GCA	ACG	ACCI	CTA	TATO	TCGI	CT C	AAGI	TCCTA	60
CTAT	CATG	TA I	GCTG	TCAC	T AC	AGAA	TAAA	TTI	TGTC	TAT	AGCT	GGCA	AG F	AAGC	CATCA	120
CATA	CATT	CT G	ATGG	TGTA	G GC	TCCA	CATO	ACA	GTAA	GCA	TTTG	TATA	AG C	GCTG <i>I</i>	ATCACA	180
TAGG						mm00		3 mc	mc a	NCC.	מממ	ጥርጥ	ጥልሮ	GC A	GAA	233
TAGG	GTGC	TA C	CGAC	CTAG	ic CA	TTGC	IAC	Met	Ser	Thr	Lvs	Ser	Tvr	Ala	Glu	
								1			-1-	5				
						- 1										
AGG	GCC	AAG	GCA	CAC	AAT	TCG	CCA	GTT	GCT	AGA	AAG	CTT	CTG	GCA	TTG	281
Arg	Ala	Lys	Ala	His	Asn	Set	Pro	Val	Ala	Arg	Lys	Leu	Leu	Ala	Leu	
	10					1/5					20					
		63.6	220	777	NCC.	AAT	רייר ר	ጥርር	ር ር ጥ	тсс	СТТ	GAT	GTG	CGG	ACG	329
ATG	CAC	GAG	T.ve	AAA T.vs	Thr	Asn	Leu	Cvs	Ala	Ser	Leu	Asp	Val	Arg	Thr	
25	птэ	GIU	БуЗ	1 , 5	30	7		- 2		35		_			40	
						1										
TCT	AGA	AAG	CTT	CTG	GAG	CTA	GCA	GAC	ACG	CTG	GGA	CCG	CAC	ATT	TGT	377
Ser	Arg	Lys	Leu		Glu	Leu	Ala	Asp	Thr	Leu	Gly	Pro	His	11e 55	Cys	
				45	- 1				50					33		
CMC	cmc	7 7 C	A C A	$C \Lambda T$	GT/C	GAC	ልጥል	СТС	ACG	GAC	TTC	GAC	ATC	GAG	ACG	425
T.e.ii	Len	T.VS	Thr	His	Val	Asp	Ile	Leu	Thr	Asp	Phe	Asp	Ile	Glu	Thr	
neu	Deu	270	60		T		•	65		_			70			
					-											472
ACA	GTC	AAG	CCG	CTG	CAG	CAG	CTT	GCG	GCT	AAG -	CAC	AAC	TTC	ATG	ATC	473
Thr	Val		Pro	Leu	qln	Gln		Ala	Ala	Lys	His	Asn 85	Pne	мет	TTe	
		75			1		80					83				
mm <i>C</i>	CAC	CAC	CCC	אמ	I TO	CCT	GAC	Aጥጥ	GGC	AAC	ACG	GTT	AAG	CTG	CAG	521
Phe	GAG	Asp	Ara	Lvs	Phe	Ala	Asp	Ile	Gly	Asn	Thr	Val	Lys	Leu	Gln	
1 116	90	1101	9	_, 5	[95			-		100					
-					1											
TAC	TCC	TCC	GGC	GTG	TAC	CGT	ATC	GCG	GAG	TGG	GCG	GAT	ATT	ACC	AAT	569
Tyr	Ser	Ser	Gly	Val		Arg	Ile	Ala	Glu			Asp	Ile	Thr	Asn	
105					110					115					120	

							17		4								
GCA	CAC	GGC	GTC	ACC	GGC	CCC	GGT	GTG	ATA	GCC	GGG	CTG	AAG	GAG	GCT		617
Ala	His	Gly	Val		Gly	Pro	Gly	Val		Ala	Gly	Leu	Lys	Glu	Ala		
				125					₽30					135			
			GCC	503	030	C2.2	ccc	200	CCC	ጥጥር	СТС	ልጥር	СТС	GCA	GAG		665
GCG	AAA	CTG	Ala	TCA	CAG	GAA	Pro	AGG	GGG	Len	Leu	Met	Leu	Ala	Glu		
AIA	гÀг	Leu	140	Ser	GIII	GIU	FIO	145	Gry		200		150				
			140					7									
CTC	TCT	TCT	CAG	GGC	TCT	TTG	GCG	dec	GGA	GAC	TAT	ACC	GCG	GGC	GTC		713
Leu	Ser	Ser	Gln	Gly	Ser	Leu	Ala	Arg	Gly	Asp	Tyr	Thr	Ala	Gly	Val		
		155					160	1				165					
					ame	CAC	CAA	CAC	ጥጥጥ	CTC	ልጥር	GGG	ጥጥር	АТС	GCG		761
GTT	GAA	ATG	GCG Ala	AAG	CTG	Acn	GAA	Acn	Dhe	Val	Tle	Glv	Phe	Ile	Ala		, , ,
vaı	170	Met	Ala	гуз	ьеп	175	914	rab	riic	Vul	180	011					
	170					1,5					•						
CAG	CGT	GAC	ATG	GGT	GGG	CGT	G¢A	GAC	GGC	TTT	GAC	TGG	CTC	ATC	ATG		809
Gln	Arg	Asp	Met	Gly	Gly	Arg	Ala	Asp	Gly	Phe	Asp	Trp	Leu	Ile	Met		
185					190					195					200		
									003	C N C	ccc	CITIC	CCC	CAG	CAG		857
ACC	CCG	GGG	GTT Val	GGC	CTG	GAC	AGD	AAA	GGA	Acn	GGC	Len	Glv	Gln	Gln		00.
Thr	Pro	GTÄ	vaı	205	ьeu	Asp	Asp	гуз	210	АЗР	GLY	Deu	011	215	0		
				203		- 1											
TAC	CGC	ACG	GTG	GAT	GAG	GTC	GTC	AGC	GAC	GGT	ACC	GAT	GTG	ATC	ATT		905
Tyr	Arg	Thr	Val	Asp	Glu	val	Val	Ser	Asp	Gly	Thr	Asp	Val	Ile	Ile		
			220					225					230				
				ama		1	220	CCA	707	CAC	CCC	AAG	GTC	GAG	GGT		953
GTT	GGC	AGA	GGG	CTC	TTT	ASD	Tue	GGA	Ara	Asp	Pro	Lvs	Val	Glu	GGT Gly		•••
vai	GIĀ	235		теп	FIIC	Lap	240		11119	110 P		245			- 2		
						1											
GCC	CGC	TAC	CGC	AAG	GCC	GGT	TGG	GAG	GCT	TAC	TTG	CGC	CGT	ATG	GGC		1001
Ala	Arg	Tyr	Arg	Lys	Ala	Gly	Trp	Glu	Ala	Tyr	Leu	Arg	Arg	Met	Gly		
	250					255					260						
		=00	TAG	mama	mod .	cmcc	cccc	CN C	አርሞል	ጥልጥል	e ec	ссат	ጥርርል				1050
	ACT			TCTA	TCG	CTGG	CGCC	CAC	AGIA	INIA		JOHI	1001	-			
265		Ser			-												
203					-												
CCG	CCGA	TTA	CCAT	CTCA	G¢ A	ACCT	TTTT	G TA	ATTA	TATG	CCC	CTAT	TGC	CCTI	ATTTC	С	1110
					1									-0.0	m> 00>1	•	1170
GAG	CTGG	TGC	CGGG	ATCG	gr r	TATA	.GACG	iG GC	AACA	AGTT	GAT	'ACT'I	"TGT	TCAG	TAGCA'	ľ	1170
003	maa.	202	cmmc	CACC	- T	cccc	ጥርጥር	C AA		ጥሮርር	CGC	GGAT	TAAT	TCGI	ATTAC	С	1230
GCA	TCCA	ACA	CTTG	CAGG		ಲಲಲ	1010	IG AA		1000							
CGC	АСТТ	CGT	GAAG	TATI	Б С Т	TATT	GAAA	LA A	CTTC	ACTT	TGG	GCTA	ACT	AGAG	CCATA	A	1290
					l												
CTC	GACA	CAA	GCCC	CTTC	CT A	CACA	CTTC	G AG	CTGG	GACT	' AAA	AGTGA	CAA	CGAA	TAGCA	A	1350

18 ATAATTAGCA AATATGGATG CGTTGAATTQ (2) INFORMATION FOR SEQ ID NO: 2: SEQUENCE CHARACTERISTICS: (i) (A) LENGTH: 267 amino acids (B) TYPE: amino acid (D) TOPOLOGY: linear (ii) MOLECULE TYPE: protein SEQUENCE DESCRIPTION: SEQ ID NO: 2: (xi) Met Ser Thr Lys Ser Tyr Ala Glu Arg Ala Lys Ala His Asn Ser Pro 1 5 10 15 Val Ala Arg Lys Leu Leu Ala Leu Met His Glu Lys Lys Thr Asn Leu 20 25 30 Cys Ala Ser Leu Asp Val Arg Thr Ser Arg Lys Leu Leu Glu Leu Ala 40 45 35 Asp Thr Leu Gly Pro His ‡le Cys Leu Leu Lys Thr His Val Asp Ile 50 55 60 Leu Thr Asp Phe Asp Ile Glu Thr Thr Val Lys Pro Leu Gln Gln Leu 65 75 Ala Ala Lys His Asn Phe Met Ile Phe Glu Asp Arg Lys Phe Ala Asp 85 90 Ile Gly Asn Thr Val Lys Leu Gln Tyr Ser Ser Gly Val Tyr Arg Ile 105 100 Ala Glu Trp Ala Asp tle Thr Asn Ala His Gly Val Thr Gly Pro Gly 120

Arg Gly Leu Leu Met Leu Ala Glu Leu Ser Ser Gln Gly Ser Leu Ala 145

Arg Gly Asp Tyr Thr Ala Gly Val Val Glu Met Ala Lys Leu Asp Glu 165

170

175

Val Ile Ala Gly Leu/Lys Glu Ala Ala Lys Leu Ala Ser Gln Glu Pro

140

135

Asp Phe Val Ile Gly Phe Ile Ala Gln Arg Asp Met Gly Gly Arg Ala 180 185 190

Asp Gly Phe Asp Trp Leu Ile Met Thr Pro Gly Val Gly Leu Asp Asp Lys Gly Asp Gly Leu Gly Gln Gln Tyx Arg Thr Val Asp Glu Val Val Ser Asp Gly Thr Asp Val Ile Ile/Val Gly Arg Gly Leu Phe Asp Lys Gly Arg Asp Pro Lys Val Glu Sly Ala Arg Tyr Arg Lys Ala Gly Trp Glu Ala Tyr Leu Arg Arg Met Gly Glu Thr Ser